

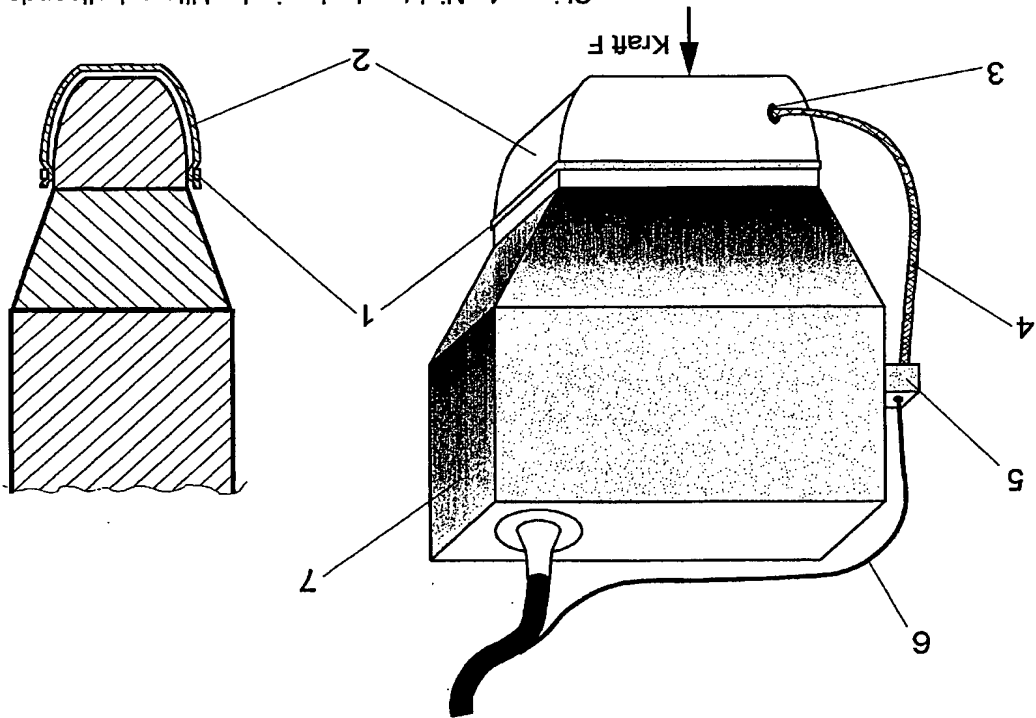


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SKIZZE 4: Nischendoskopische Ultraschallsonde



④ Erfindest:  
④ Ernest, Helmut, Prof. Dr.-Ing., 91341 Röttenbach,  
DE; Lorenz, Andreas, Dipl.-Ing., 44789 Bochum, DE;  
Wiebe, Peter, Dipl.-Ing., 58285 Gevelsberg, DE  
④ Anmelder:  
④ gleich Anmelder  
④ Erfindest:  
④ gleich Anmelder

Die folgenden Angaben sind den vom Anmelder eingerichteten Unterlagen entnommen  
④ Ein sonographisches Elastographiesystem

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## Beschreibung

Die Erfahrung betrifft eine Meßrichtung entsprechend dem Oberbegriff des Anspruchs 1.

## Anwendungsbereich

Die mechanischen Eigenschaften von biologischem Gewebe (z. B. Elastizitätsparameter) sind für die Beurteilung des Zustandes des Gewebes von großem Interesse. In der medizinischen Diagnostik deuten Veränderungen der Elastizitätseigenschaften auf histologische und u. U. pathologische Veränderungen hin. Allgemein bekannt sind Prozesse wie die Bildung von Geschwülsten und Verkürzungen ('Knoten'), die z. B. manuell tastbar sind. In der Landwirtschaft ist für die Beurteilung der Qualität des Fleisches von Schlachtrind die Kenntnis mechanischer Gewebeigenschaften ebenfalls von Interesse.

## Stand der Technik

Die sogenannte Taubflöpfung ist ungenau und unempfindlich. Wesentlich besser ist in dieser Beziehung die sogenannte Elastographie, bei der elastische (Gewebe)geschwindigkeiten technisch erfaßt und z. B. in Form von Schmittintervallen quantitativ oder quantitativ visualisiert werden. Dabei benötigt man sich hauptsächlich des Ultraschalls, wie er als bildgebendes Verfahren in der medizinischen Diagnostik eingesetzt wird. In zentral nachmehrander aufgenommenen Schallfeldern können geringe Verschiebungen oder Verformungen innerhalb der dargestellten Gewebestruktur durch Auswertung der Bildsequenzen erfaßt und ausgewertet werden. Wird auf einen Gewebebereich ein mechanischer Druck ausgeübt, der eine Verformung des Gewebes zur Folge hat so verformen sich Bereiche mit unterschiedlichen Elastizitätseigenschaften verschiedenartig. Das Elastographiesystem wertet diese Verformungen durch den numerischen Vergleich der Elastizitätsparameter aus und stellt die unterschiedlichen Elastizitätsparameter im Bild dar. Die nun erfolgende Kompression des Gewebes, die z. B. extern provoziert wird, ist nur gering und begünstigt bei Anwendung des üblichen diagnostischen Ultraschalls von Millimetern. Wichtig dabei ist eine quantitative Kontrolle des Kompressionsmaßnahmen.

Ein Verfahren der Ultraschall-Elastographie von Körpergewebe ist erstmalig in einem Aufsatz von J. Opipri et al im Jahre 1991 [1], [2] beschrieben worden. Dabei werden Ultraschallbilder bzw. die entsprechenden hochfrequenten Ultraschalltechnologien so ausgewertet, daß Verschiebungen des Körpergewebes zwischen zwei aufgenommenen Kompressionsaufnahmen Gewebehüften berechnet werden. Auf diese Weise lassen sich die bereits erläuterten Rückschlüsse auf die Elastizität des Organs bis hin zu einer quantitativen Abbildung des Elastizitätsmoduls erzielen.

In der Literatur werden verschiedene Ansätze vorgestellt, mit denen die Abbildungseigenschaften eines Elastographie-Systems verbessert werden können. Versprechend im Hinblick auf das Signal-Rauschverhältnis und den Kontrast der Elastographiebilder sind Ansätze, die mehrfache Kompressionsstufen das abzuhörenden Gewebe auswerten. So wird in diesem Ansatz bis zu 120 Bilder in einer Aufnahmeserie aufgenommen und mit bekannten Methoden der Ultraschall-Elastographie [3], [4] weiterverarbeitet.

## Nachteile des Standes der Technik

Ein Nachteil der bisher publizierten bzw. praktizierten Verfahren ist, daß zu keiner Zeit die Kraft auf das Gewebe

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## Beschreibung

bekannt ist, die wertvolle Informationen bei einer quantitativen Rekonstruktion des Elastizitätsmoduls liefern kann. Außerdem ist ein solcher Ansatz speziell, rechner- und damit kostenintensiv, da Auswahlkriterien dazu herangezogen werden müssen, optimale Bildfolgen aus den aufgezeichneten Daten auszuwählen und weiterzuarbeiten.

## Aufgabe der Erfindung

Die Aufgabe der Erfindung ist es, ein Ultraschallbild der in vorgegebenen Kompressionsstufen aufzunehmen, um eine Anzeige der elastischen Eigenschaften von Körpergewebe effizienter zu machen und durch genaue Messung und Kontrolle des applizierten Drucks eine quantitative Rekonstruktion des Elastizitätsmoduls zu unterstützen.

## Lösung der Aufgabe

Diese Aufgabe wird durch ein McSystem mit den Merkmalen des Anspruchs 1 gelöst.

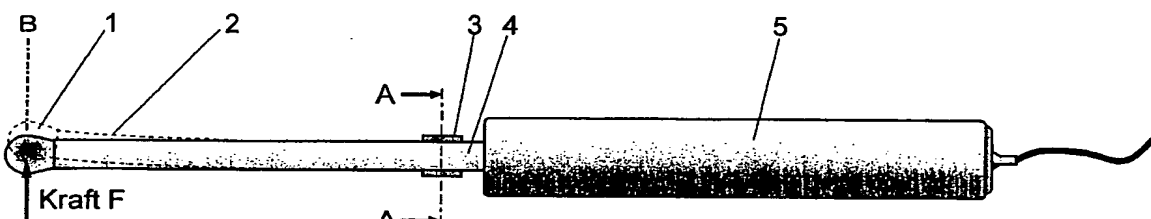
Die Kompression des Gewebes wird durch den Ultraschallwandler, mit dem die Ultraschallbilder aufgenommen werden, hervorgerufen. Die Kraft, die der Schallwandler auf die Gewebe ausübt, wird durch eine mit diesem Wandler verbundene Vorrichtung gemessen, und es werden Ultraschallbilder in vorgegebenen Kompressionsstufen aufgenommen. Insbesondere soll sich bei endoskopischen Schallwänden, (z.B. bei einer transstalakalen oder transvaginalen Sonde), die für gesonderte Kraftmeßvorrichtungen nur wenig Platz erhalten, die Kraft mittels eines am Schallkopf angebrachten Dehnungssensors nach dem Blaschko-Konzept bestimmen. Für nicht-endoskopische Schallköpfe, d.h. Schallköpfe die auf der Körperebene appliziert werden, kann eine wassergetüpfelte, schalldurchlässige Koppe auf dem Schallkopf angebracht werden. Eine Kraftmessung ist dann mittels eines hydrostatischen Drucksensors möglich.

## Vorteile der Erfindung

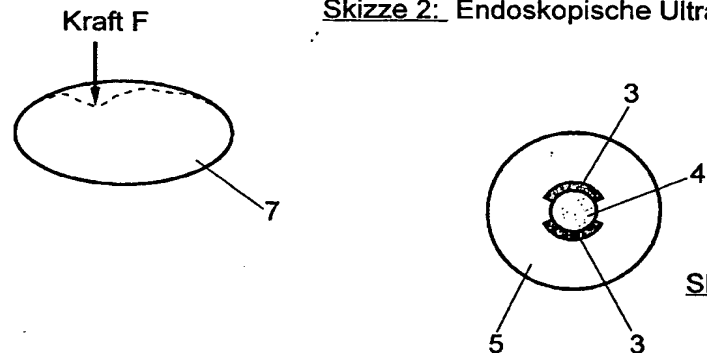
Dieser Ansatz erlaubt eine Montage der Sensoren auf handhabbare Ultraschallsonden mit nur geringfügiger Modifikation der Schallkopfe. Es werden keine geschäftsähnlichen Verhinderungen am Ultraschallgerät vorgenommen. Weiterhin erfolgt die Bildaufnahme zu gezielt vorgegebenen Kraft- bzw. Drucksäulen, was den Speicher- und Rechenaufwand im Vergleich zu bekannten Ansätzen erheblich reduziert. Die Erfindung macht bei der Anwendung starker endoskopischer Schallsonden eine quantitative Elastographie erst möglich.

## Literaturangaben

- [1] Opipri J., Céspedes I., Ponnakkani H., Yazdi Y., Li X.: Elastography: A quantitative method for imaging the elasticity of biological tissues. Ultrasonic Imaging 13, 111-114, 1991.
- [2] Céspedes I., Opipri J., Ponnakkani H., Maklad N.: Elastography: Elasticity imaging using ultrasound with application to muscle and breast imaging in vivo. Ultrasonic Imaging 15, 71-88, 1993.
- [3] O'Donnell M., Slovoda A. R., Shapo B. M., Emilia nov S.: Internal displacement and strain imaging using ultrasonic speckle tracking. IEEE transactions on ultrasonics, ferroelectrics and frequency control, 41, 314-322, May 1994.
- [4] Lubinski M. A., Emiliyanov S. Y., Raghavan K. R.: Internal displacement estimation using tissue incompressibility



Skizze 2: Endoskopische Ultraschallsonde (schematisch)



Skizze 3: Schnitt A-A zu Skizze 2

Nummer:  
Int. Cl. 6:

Offenlegungstag:

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mit handelsüblichen Ultraschallgeräten bzw. Sanden mit geringfügiger Modifikation der Schallsonde erfüllt werden kann, ohne gerichtsbauliche Veränderungen am Ultraschallgerät vornehmen zu müssen.

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11/2x 1 Seite(n) Zeichnungen

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tor, not being sensitive to phase changes, would still be more robust than the RFT crosscorrelation estimators. In fact, phantom experiments were used to show that the centroid estimator could generate quality elastograms at applied strains as high as 3%. While the cross-correlation based elastograms are extremely noisy, furthermore, simulation results showed that the spectral centroid shift method provided a jitter insensitive method of estimating the strain. Therefore, spectral strain estimation may be particularly useful for obtaining good elastograms in noisy jitter environments produced by unpredictable tissue and/or system motion. This may constitute a major advantage, since elastography might be practiced using the same clinical guidelines employed by ultrasound; i.e., using a hand-held transducer. In addition, the jitter resistance of the centroid estimator could make it suitable for use in intravascular elastography *in vivo*, a task that has not been demonstrated as feasible using cross-correlation techniques. Future investigations will involve theoretical study of the performance of spectral (*i.e.*, frequency and shift) and bandwidth strain estimators as well as experimental verification of their jitter insensitivity.

[0083] The foregoing disclosure and description of the invention are illustrative and explanatory. Various changes in the size, shape, and materials, as well as in the details of the illustrative construction may be made without departing from the spirit of the invention.

What is claimed:

1. A method for measuring strain in a target body comprising:
  - a. acoustically coupling a transducer to the outer surface of a target body such that the path of a beam emitted from the transducer defines a transducer axis;
  - b. emitting first a pulse of ultrasound energy into the target body along the transducer axis;
- c. receiving a first reflected signal with the transducer;
- d. storing the first reflected signal;
- e. allowing the target to change dimensions along the axis defined by the transducer;
- f. emitting a second pulse of ultrasound energy into the target body along the transducer axis;
- g. receiving a second reflected signal with the transducer;
- h. storing the second reflected signal;
- i. selecting a portion of the first and second reflected signals;
- j. computing the frequency spectrum of each of the selected portions of the first and second selected signals;
- k. computing the shift between the computed spectra; and
- l. normalizing the computed shift to one of the computed spectra.

2. The method of claim 1 wherein allowing the target to change dimensions is accomplished by applying a compressive force to the target.
3. The method of claim 1 wherein allowing the target to change dimensions is accomplished by reducing a compressive force to the target.
4. A method of claim 1 wherein computing the frequency spectrum of each of the selected portions of the first and second selected signals is accomplished using Fourier analysis.

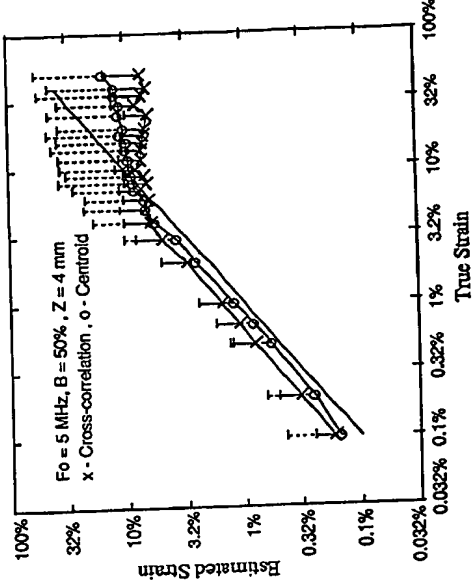
\* \* \* \* \*

(54) POWER SPECTRAL STRAIN ESTIMATORS IN ELASTOGRAPHY (57) ABSTRACT

Elastography can produce quality strain images *in vivo* and *in vitro*. Standard elastography uses a coherent cross-correlation technique to estimate tissue displacement and tissue strain using a subsequent gradient operator. While coherent estimation methods generally have the advantage of being highly accurate and precise, even relatively small undesired motions are likely to cause enough signal decorrelation to produce significant degradation of the elastogram. For elastography to become more universally practical in such applications as hand-held, intravascular and abdominal imaging, the limitations associated with coherent strain estimation methods that require tissue and system stability, must be overcome. In this paper, we propose the use of a spectral shift method that uses a centroid shift estimate to measure local strain directly. Furthermore, we also show theoretically that a spectral bandwidth method can also provide a direct strain estimation. We demonstrate that strain estimation using the spectral shift technique is moderately precise, but far more robust than the cross-correlation method. A theoretical analysis as well as simulations and experimental results are used to illustrate the properties associated with this method.

(76) Inventors: Elisa Konofagou, Boston, MA (US); Jonathan Ophir, Houston, TX (US)  
Correspondence Address:  
Richard T. Redon  
Duane, Morris & Heckscher LLP  
Suite 500  
One Greenway Plaza  
Houston, TX 77046 (US)

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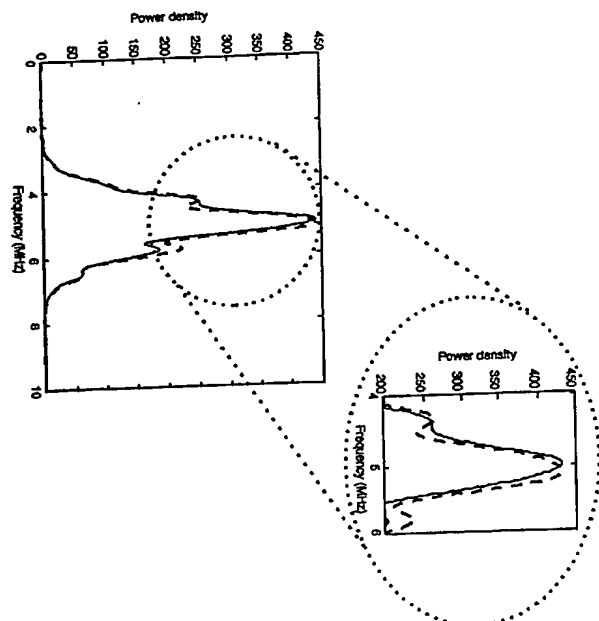


Figure 1

[6073] The strain estimation noise performance of the centroid estimator was compared to that of the standard catastrophic cross-correlation-based strain estimator with multi motion compensation, i.e., global stretching. As explained in the introduction, motion compensation was not used, since the ratio to which the robustness of the two estimators is tested is precisely the one due to axial motion bandwidth and 100 A-lines.

stimulate slip boundary conditions and was free on both lateral and elevational sides.

[A80] Two major differences can be observed between signals are completely decorrelated producing (FIG. 7a (ii))

elastogram generated using the cross-correlation algorithm provides the closest correspondence to the ideal elastogram. On the other hand, for larger applied strains (5% and 10%), the cross-correlation algorithm fails to accurately estimate tissue strain due to the increased signal-to-decoration errors (FIG. 6*c* and 6*d*, respectively). In fact, at 5% applied compression part of the inclusion is visible, being three times harder than the background and, thus, experiencing a much lower strain allowing it to be depicted with a good signal-to-noise ratio. On the other hand, the elastogram generated using the spectral centroid method at 5% and 10% compression (FIG. 6*c* and 6*d*, respectively) illustrates the robustness associated with the centroid method.

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[0075] In the next section, we present elastograms obtained using an elastographic experimental phantom. The experimental results provide a complex 3-D situation where axial, lateral and deviatoric signal decorrelation are present, unlike the 1-D situation illustrated in this section.

[0081] The new concept described in this patent is based on the direct estimation of tissue strain from the relative frequency shift in the power spectrum. The estimator becomes presented, namely the centroid shift estimator, increases its shift by calculating the relative centroid shift resulting from the applied compression. This estimator has three major characteristics: it is a) direct and b) self-centered, i.e., operates the frequency domain. The direct strain estimation assures

[0077] The ultrasound system used for taking the data was a Sonix-SCI (Sonix, Inc., San Clara, Calif.) operating with dynamic receive focusing and a single transmit focus centered at a depth of 5 cm. The transducer used was a 5-MHz linear array (40 mm) with a 60% fractional bandwidth. The digitizer used is a 8-bit digitizer (Ultrav Corp., Spring Valley, N.Y.) with a sample rate of 40 MHz. The digitized data was collected from a 40x40 mm ROI consisting of 100 A-lines (spaced at a depth of 5 mm under the transducer) centered around the transcutaneous focus. The system also included a motion controller, system, and a compression device. A personal computer controlled the operation of the entire system.

**(b) 1-D**: In order to study the performance of the centroid estimator, we used a 1-D simulation model that allowed it to scatter to move solely in the axial direction. Preliminary results obtained with these 1-D simulations are used to demonstrate the robustness of the proposed method. Since estimates as high as 10% are produced at a relatively high signal-to-noise ratio while the standard crosscorrelation-based diagnostic method practically failed beyond 1% levels of noise, the 1-D example was preferred to 2-D or 3-D simulations so that the performance of the method could be characterized independent of noise due to 2-D or 3-D motion. If the 2-D or 3-D scenario were used, the estimator would fail in lower studies, but the spectral centroid estimat-

4 mm and the size of postcompression window was changed with strain in order to assure that the same tissue information was incorporated in both the pre- and postcompressed windows. The length of the data segment incurred the usual tradeoff in spectrum estimation. A larger window length improved the spectrum as long as the data was stationary. Moreover, it was recently shown that the overlap has a more significant impact on resolution than the window length. As a result, larger windows with high overlap can generally provide both a smoothing effect on the noise and high resolution for both the time-domain and spectral estimators. This effect, however, needs to be further investigated.

[0066] The power spectra calculation of the pre- and postcompressed RF segments was performed using a 25-point frequency smoothing window unless otherwise stated. The 25 point frequency smoothing window represented only 0.6% of the entire FFR and is a relative small window. Frequency smoothing is similar to using a moving-average window, however, the averaging is performed on the complex spectrum to obtain an estimate of the power spectrum. Frequency smoothening allowed the use of a single pair of A-lines, similar to the strain estimator performed using the cross-correlation-based strain estimator. We use a 1024 point chip Z-Transform to compute the spectrum we which correspond to a 4096 point FFT (since we use only one half of the spectrum, and only the region with a sufficient signal). The mean and standard deviation of the strain estimates were obtained by processing pre- and postcompression signals with a total length of 30 mm. The corresponding SNR (ratio of the mean of the estimated strain to its standard deviation) values were obtained using Monte Carlo simulations with 25 independent realizations for each strain value. The simulated Strain Filters were obtained by plotting the SNR estimates for the whole range of the applied tissue strains. The Strain Filter, typically addresses the limitations of the ultrasound system (such as time-bandwidth product, center frequency and sonographic SHR) as well as the signal processing algorithms used to process the signals through the introduction of constraints in the available elastographic SNR, resolution, sensitivity and strain dynamic range.

[0067] The robustness of the strain estimations was also evaluated by introducing jitter errors in the scatterer positions before generating the post-compression signals. The jitter in the scatterer positions followed a normal distribution that varied randomly from zero to the maximum value of the jitter introduced. For the larger jitter values the scatterers could move out of the window of estimation. The strain estimation accuracy and precision for the coherent estimations were expected to deteriorate under these conditions since they depend on the relative motion of the scatterers themselves with compression. However, the centroid estimator, being incoherent, was expected to show strain estimation with a reasonable SNR, even at high jitter levels.

[0068] An example of the frequency shift on simulated spectra of entire A-lines (40 mm in length) is shown in FIG. 1 for the case of 1% applied strain. Comparison of the strain estimators using the coherent cross-correlation and centroid based algorithms are presented in FIGS. 2 and 3 for the 1-D simulations. The mean strain estimates and their standard deviation are presented in FIG. 2 and the respective simulated strain filters are presented in FIG. 3. The results in FIG. 2 illustrate that the strain estimates from both estima-

tors follow the theoretical curve (straight solid line at 45 degrees) for strains less than 5%, where the cross-correlation strain estimator begins to level off, and at 9% where the centroid strain estimator crosses the theoretical curve. Both estimators are biased with a small overestimation of the strain seen for strains lower than 5%. For larger strains, the centroid estimator underestimates the actual strain values with a larger bias in the estimated strain value. This bias in the strain estimator for the centroid estimator is at least partly due to the bandwidth broadening, as discussed in the theory section. The bias in the cross-correlation based strain window that corrupt the time-delay estimates. These bias errors can be reduced by temporally stretching the post-compression data. Overall, when compared to the standard elastographic coherent estimator, the centroid strain estimator provides a biased but more robust strain estimate.

[0069] In FIG. 3, the simulation Strain Filters for the two algorithms illustrate the noise performance of the estimators. Note that the coherent cross-correlation strain estimator provides accurate and precise strain estimates for strains less than 2%, since the variance decreases rapidly beyond this strain value; however, for larger strains the performance degrades significantly. On the other hand, the centroid strain estimator, not being sensitive to phase, provides a robust strain estimate even at very large strains close to 30 %. The SF for the centroid strain estimator indicates a reasonable SNR for low tissue strains as well as an increase in the SNR, observed for larger strains where the cross-correlation strain estimator is limited by signal decorrelation errors. Due to its lower precision the centroid estimator works best at large strains, where the strain is greater and therefore the signal-to-noise ratio (assuming the variance remains constant) increases, as shown in FIG. 3. The simulations therefore show the robustness of the spectral centroid strain estimator to large applied strains and increased jitter, errors that are most likely to be encountered in hand-held or intravascular or abdominal elastography.

[0070] Next, we investigated the sensitivity of the cross-correlation and spectral strain estimator to variations in the scatterer positions caused due to axial jitter. The results are illustrated in FIGS. 4 and 5 for the cross-correlation and centroid strain estimators respectively. Note that the coherent cross-correlation-based strain estimator is more susceptible to jitter than the centroid strain estimator. The noise performance of the cross-correlation estimator drops by about 50% with an increase in the maximum value of the jitter by 50 ns. However, in the case of the centroid strain estimator the noise performance remains at the same high level, even at jitter magnitudes of 100 ns. The simulations, therefore, show the robustness of the spectral centroid strain estimator to large applied strains and increased jitter, errors that are most likely to be encountered in hand held or intravascular or abdominal elastography. The two following sections compare elastograms obtained with these two estimators in the case of a 1-D finite-element simulation and an experimental phantom.

#### [0071] Elastograms using simulated data

[0072] After testing the properties of the new estimator in the previous section, elastograms were generated for a simulated single inclusion phantom under uniform compression. For the calculation of the displacements, we used a

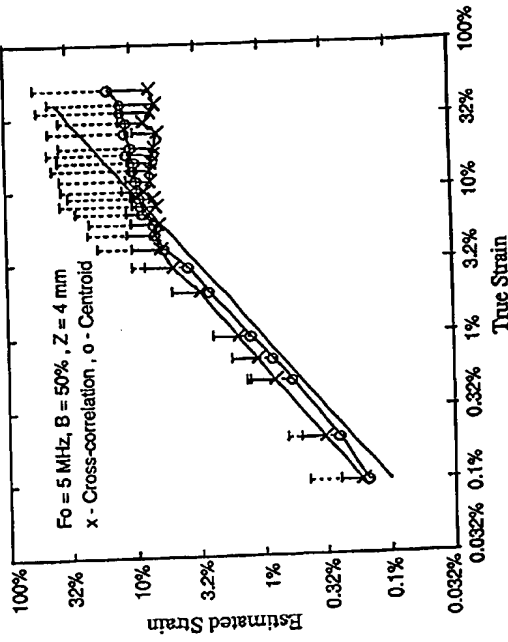


Figure 2

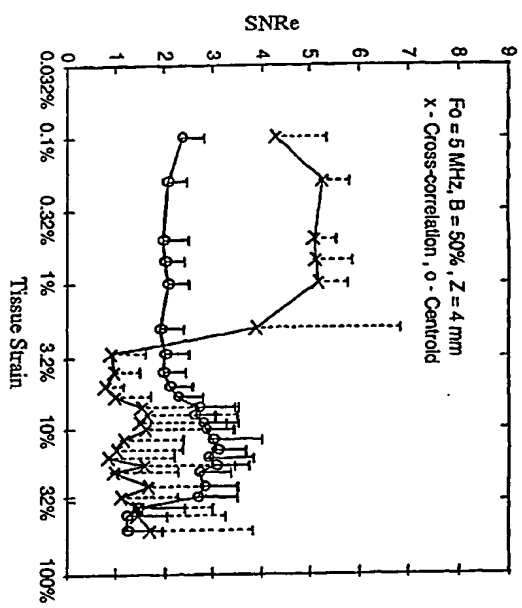


Figure 3

the centroid estimator. Otherwise, if  $B_0$  is infinite and/or if  $k$  is zero, constant  $A$  in Eq. (16) will always be zero regardless the strain.

[0043] For relatively large strains, using the more general form of Eq. (3), Eq. (16) becomes

$$\frac{f_{2t} - f_{1t}}{f_{1t}} = \frac{1}{1-s}$$

[0044] or, by solving for the strain,

$$s = \frac{f_{2t} - f_{1t}}{(\lambda - 1)f_{1t} + f_{2t}} \quad (19)$$

[0045] which is a less straightforward, but still a direct way of estimating higher strains.

[0046] Eq. (16) is reminiscent of the well-known Doppler effect according to which the ratio of the centroid shift to the center frequency may provide a reliable measure of velocity.

According to Eq. (16), in the case of strain, a similar effect also occurs. Also, among others have shown how in broadband Doppler the bandwidth of the resulting spectrum also changes with velocity and that the output RF Doppler spectrum is a frequency-shifted and compressed (or stretched) replica of the transduced one. Similarly, in the case of strain measurement, that the following expression provides a direct estimation of strain:

$$\frac{f_{2t} - f_{1t}}{f_{1t}} = ts \quad (20)$$

[0047] where  $t$  are the post- and precompression bandwidths, respectively and  $k$  is constant. So, strain, like velocity, introduces these two coupled effects of centroid shift and bandwidth variation in the power spectrum. Spectral broadening (i.e., in the case of compression) or contraction (i.e., in the case of tension) and (over) compression of the spectrum (i.e., in the case of compression) or contraction frequencies separated by a shift,  $M$ , and a bias term  $B$  denoting the spectral broadening (on compression) due to strain  $s$ :

$$f_{2t} - f_{1t} = tsM + B \quad (21)$$

[0048] where

$$B = B_0 + B_1s - B_2 \quad (22)$$

[0049] In order to estimate the strain without the bias associated with spectral broadening, the following equation can be used that results from Eqs. (19), (20), (21) and (22):

$$ts = \frac{B_0 + B_1s - B_2}{M} \quad (23)$$

[0050] and solving for strain, the unbiased estimator is given by

$$s = \frac{B_0 + B_1s - B_2}{Mf_{1t} - Mf_{2t}} \quad (24)$$

[0051] However, Eq. (24) requires a bandwidth estimation and since the bandwidth estimator is not part of this study, we use Eq. (16) as the strain estimator and show a bias with simulations, which is partly due to the previously described bias due to spectral broadening.

#### DESCRIPTION OF THE DRAWINGS

[0052] FIG. 1 is a graph of power density versus frequency.

[0053] FIG. 2 is a graph of estimated strain versus true strain.

[0054] FIG. 3 is a graph of signal to noise ratio versus tissue strain.

[0055] FIG. 4 is a graph of signal to noise ratio versus tissue strain.

[0056] FIG. 5 is a graph of signal to noise ratio versus tissue strain.

[0057] FIG. 6 (i)-(c) are elastographs.

[0058] FIG. 7 (i)-(c) are elastographs.

[0059] FIG. 8 (i)-(c) are elastographs.

[0060] FIG. 9 comprises various elastographs.

[0061] FIG. 10 is a graph of power spectrum versus frequency.

[0062] 4. Description of the Preferred Embodiments

[0063] Simulation results using a 1-D scattering model is used in this section to illustrate the performance of the centroid strain estimator. Strain estimation using the centroid estimator is also compared to the standard cross-correlation based algorithm.

[0064] Monte-Carlo simulations in MATLAB (Mathworks, Inc., Natick, Mass., USA) are used to generate pre- and post-compression RF signals for a 30 mm target segment and sampled at 48 MHz. The speed of sound in tissue was assumed to be constant at 1540 ms. The PSF was simulated using a Gaussian modulated cosine pulse with a wave number=0.04 mm (5 MHz center frequency, 50% bandwidth), and a 0.2138 mm standard deviation unless stated otherwise. The scattering function consisted of randomly distributed point scatterers following a uniform distribution with density of 40 scatterers/pulse-width in order to simulate Gaussian statistics. We assume that the uniformly distributed scatterers are of sufficient number to generate an echo signal with circular Gaussian statistics. The PSF was convolved with the scattering function to obtain the pre-compression RF signal. The post-compression signals were generated after applying a uniform compression of the point scatterers, and convolving the compressed point scatterers with the original PSF.

[0065] Spectral strain estimation (following Eqs. 16 and 11) was performed using pre- and postcompressed power spectra of windowed RF signals. The signal length equaled

similar result is found later (Eq. 16) using the spectrum of the received signal. In the section below we use Eq. (10) as a guide in the formulation of the new estimator.

[0027] Effect of strain on the spectrum of the received signal

[0028] The centroid of the power spectrum of the received signal is defined as follows:

$$f_c = \frac{c}{2\pi} \int_{-\infty}^{\infty} R_s(\lambda) d\lambda \quad (11)$$

[0029] The centroid estimate for the precompression power spectrum is given by:

$$f_{c0} = \frac{c}{2\pi} \frac{k_0^2 l_0^2 + k_0^2}{l_0^2 + k_0^2} \quad (12)$$

[0030] In a similar manner we can derive the expression for the centroid of the post-compression power spectrum by replacing  $k_0$  and  $l_0$  in Eq. (9) by their corresponding parameters in the post-compression power spectrum, i.e.,  $k_A$  and  $l_A$ .

[0031] (as indicated from Eq. 7) we obtain:

$$\frac{l_A}{a} = \frac{c}{2\pi} \frac{k_A^2 l_A^2 + k_A^2}{l_A^2 + k_A^2} \quad (13)$$

[0032] Note that the PSF parameters remain unchanged. Since both centroids depend on the center frequencies and bandwidths of the scattering function and the PSF and by consulting Eq. (10), we normalize this effect by using the following ratio as a candidate strain estimator:

$$\frac{f_A - f_{c0}}{f_{c0}} = \frac{\frac{c}{2\pi} \frac{k_A^2 l_A^2 + k_A^2}{l_A^2 + k_A^2} - \frac{c}{2\pi} \frac{k_0^2 l_0^2 + k_0^2}{l_0^2 + k_0^2}}{\frac{c}{2\pi} \frac{k_0^2 l_0^2 + k_0^2}{l_0^2 + k_0^2}} \quad (14)$$

[0033] The parameters  $l_0$  and  $l_A$  are related to  $B_0$  and  $B_A$ , the equivalent noise spectral bandwidths for the scattering and PSF spectra, through

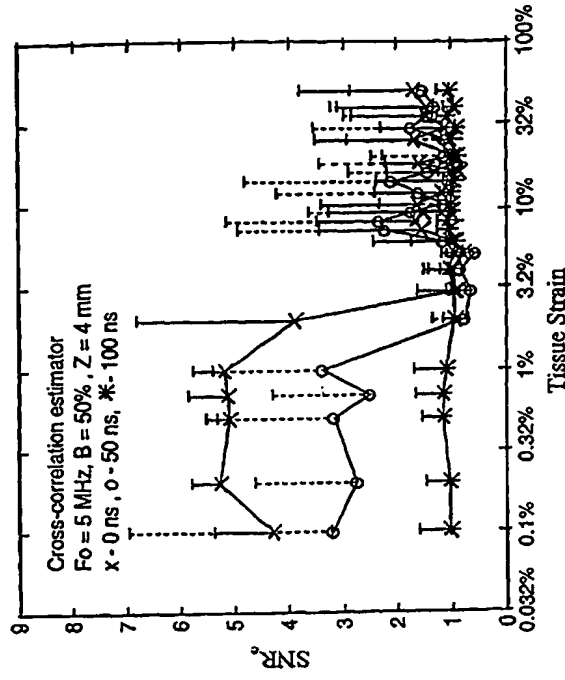


Figure 4

$$R_0 = \frac{1}{2\pi f L_0} \quad \text{and} \quad f_0 = \frac{1}{2\pi f L_0}$$

[0034] respectively.

[0035] However, the PSF bandwidth is typically much smaller than the bandwidth of the scattering function, i.e.,  $B_0 \gg B_s$ ; thereby,  $L_0 \ll L_s$  and therefore,  $a^2 L_s^2 \gg a^2$ . After cancellation of common terms in the numerator and denominator of Eq. (14), we obtain

$$\frac{f_A - f_{c0}}{f_{c0}} = \frac{(a-1) \frac{k_A^2}{k_0^2} \frac{l_A^2}{l_0^2}}{\frac{k_A^2}{k_0^2} + \frac{l_A^2}{l_0^2}} \quad (15)$$

[0036] or, from the small strain approximation case of Eq. (3) (i.e., in mathematical terms, for strains less than 10%),

$$\frac{f_A - f_{c0}}{f_{c0}} \approx \frac{f_A - f_{c0}}{f_{c0}} \quad (16)$$

[0037] where  $A$  is given by

$$A = \frac{1}{\frac{k_A^2}{k_0^2} + \frac{l_A^2}{l_0^2}} \quad (17)$$

[0038] Inspection of Eq. (16) leads to the following interesting observations:

[0039] The relative spectral centroid shift can be used as a direct strain estimator. We can also observe a direct analogy between the classic definition of strain (Eq. (4)) and the estimator of Eq. (16), which establishes this method as a simple and straightforward way of estimating the strain.

[0040] When the strain is positive (or, compressive), a frequency upshift occurs, i.e.,  $f_A > f_{c0}$ . Conversely, a tensile (or, negative) strain results in a frequency downshift, i.e.,  $f_A < f_{c0}$ . Therefore, the estimator of Eq. (16) provides directly not only the magnitude of the strain but also its sign.

[0041] Since constant  $A$  is independent of the strain, it will introduce a uniform bias on the resulting elastogram. This should not affect the resulting elastogram, since the latter depicts relative values of strain. The reader should note that the effect of local bandwidth variations is ignored.

[0042] The scattering spectrum must be a bandpass and bandlimited spectrum in order to estimate the strain using

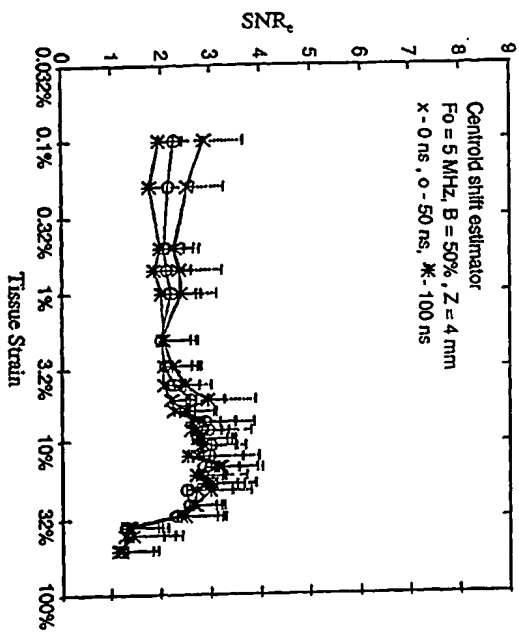


Figure 5

tions that deal with the development of alternative strain estimators as well as bandwidth estimations will not be reported in this study.

[0012] The spectral centroid has been widely used in estimating the Doppler shift, attenuation and backscattering. The theory underlying the use of centroid strain estimators is presented in the next section. One-dimensional (1-D), instead of two-dimensional (2-D), motion simulations are used in order to more accurately study the performance of the estimator, i.e., independent of the effect of signal decorrelation in two dimensions that complicates the measurements. Simulation results in 1-D illustrate the insensitivity of the centroid strain estimator to signal decorrelation effects. It is important to note that, as mentioned earlier, decorrelation can be due to several sources. For the purpose of this paper, we consider solely the axial decorrelation effect in this 1-D model. We, thus, assume that the robustness demonstrated by the spectral estimator vis-à-vis this effect is a more general property that can be further applied to other decorrelation scenarios. For example, it is shown in the results section how the spectral method is indeed more immune to jitter, another source of decorrelation. The class experiments illustrate the robustness of the spectral centroid estimator. The properties of the new estimator are discussed and summarized in the Conclusion section.

[0013] In this section, we show analytically that for Gaussian echo spectra, the relative spectral shift is a direct measure of tissue strain. We also show the relative bandwidth variation can also be used as a direct strain estimator.

#### [0014] Signal and noise model.

[0015] The pre- and post-compression echo signals are given as follows:

$$\begin{aligned} r_1(t) &= \alpha c(t) e^{j\omega_0 t} n_1(t) \\ r_2(t) &= \alpha c(t) e^{j\omega_0 t} n_2(t) \end{aligned} \quad (1)$$

[0016] where  $t$  is spatial variable,  $c(t)$  and  $n(t)$  are the received RF signals before and after compression, respectively,  $\alpha$  is the impulse response of the ultrasound system or point-spread function (PSF),  $c(t)$  is the scattering function,  $\alpha$  and  $n$  are independent zero-mean white noise sources and  $\alpha$  is the compression coefficient (or, strain factor) linked to strain  $s$  through

$$\alpha = \frac{1}{1-s} \approx 1+s \quad (2)$$

[0017] The approximation holds for  $s < 1$ , where the strain  $s$  for a one-dimensional homogeneous target is typically defined in mechanics by

$$s = \frac{\ell_s - \ell_0}{\ell_0} \quad (4)$$

[0023] where  $\ell_0$  and  $\ell_s$  are the pre- and postcompressed axial dimensions of the target. From Eq. (4) the reader should note that positive strain denotes compression (and  $s > 1$ ) while negative strain denotes tension (and  $s < 1$ ). The

reader should note that throughout this paper the subscripts  $1$  and  $2$  denote pre- and postcompression parameters, respectively.

[0019] Assuming that  $b(z)$  and  $c(z)$  in Eqs. (1) and (2) can be described by their autocorrelation functions that may be modeled by modulated Gaussian functions, we obtain

$$R_{11}(k) = \frac{1}{\sqrt{L_1 L_2}} \exp(-k^2 / (4L_1^2 + 4L_2^2)) \quad (5)$$

$$R_{22}(k) = \frac{1}{\sqrt{L_1 L_2}} \exp(-k^2 / (2L_1^2 + 2L_2^2)) \quad (6)$$

[0020] and

[0021] where  $L_1$  and  $L_2$  are the resolution lengths of the PSF and of the scattering function, respectively,  $k_1$  is the central spatial frequency of the PSF, and  $k_2$  is the central spatial frequency of the scattering function.

[0022] The one-sided power spectra of the pre- and post-compression RF signals (positive frequencies) are given respectively by

$$R_{11}(k) = \frac{1}{4} \exp\left(\frac{1}{2}[(k-k_1)^2 L_2^2 + (k+k_1)^2 L_2^2]\right) * N(k) \quad (7)$$

$$R_{22}(k) = \frac{1}{4} \exp\left(\frac{1}{2}[(k-k_2)^2 L_1^2 + (k+k_2)^2 L_1^2]\right) * N(k) \quad (8)$$

[0023] where  $N(k)$  and  $N(k)$  are independent power spectra of zero-mean white noise processes, i.e.,

$$\langle N^{(1)} \rangle = \langle N^{(2)} \rangle = 0$$

[0024] A brief observation of Eqs. (7) and (8) reveals the centroid shift in the scattering spectrum resulting from the compression. In other words, if  $\ell_1$  and  $\ell_2$  are the center frequencies of the scattering spectrum before and after compression, respectively, and assuming that the speed of sound in the tissue remains constant, from Eqs. (7) and (8) we have

$$\begin{aligned} f_{\ell_2} - f_{\ell_1} &= \frac{c}{2L_1} (\ell_1 - \ell_2) \\ &= (s-1)f_{\ell_1} \end{aligned} \quad (9)$$

[0025] So, the relative centroid shift in the scattering function spectrum constitutes a direct strain estimator. A

## POWER SPECTRAL STRAIN ESTIMATORS IN ELASTOGRAPHY

### BACKGROUND OF THE INVENTION

#### [0001] 1. Field of the Invention

[0002] The present invention relates to a method of measuring strain in a target body, using the transmission, reception, processing, and normalization of ultrasound signals.

#### [0003] 2. Description of the Prior Art

[0004] Imaging of elastic parameters of soft tissue has developed into a new tool for diagnosis of disease. Current estimators of tissue motion, include a time-domain cross-correlation based speckle tracking algorithm, and a Fourier based speckle phase-tracking technique. These techniques are coherent estimation techniques, i.e., these methods are sensitive to phase variations. The coherent estimation techniques generally have the advantage of being highly precise. Strain Filter (SF) analysis has shown, however, that they are not very robust in the presence of even a small amount of de-correlation between the pre- and post-compression signals. The term robustness has been used in statistical analysis to denote the good performance of statistical tests, i.e., the homogeneity of the variance calculation, even if the data deviates from the theoretical requirements. By equivalence in elastography, robustness denotes the consistently good performance of the estimator even at a high de-correlation noise level (i.e., keeping the variance of estimation at a relatively constant and low level at a large range of noise levels).

[0005] The term de-correlation as used herein, is defined as the loss of full correlation between the pre- and post-compressed windowed signal segments. Therefore, de-correlation may be encountered due to many sources, such as intrawindow motion (unrelated lateral or elevational motion, jitter (i.e., any cause of misregistration between the pre- and post-compressed A-line segments), unstable mechanical setup, etc.). The main idea in this study is to introduce a new estimator that is more immune to de-correlation compared to other estimators.

#### [0006] 3. Summary of the Invention

[0007] The tissue strain estimator is a spectral estimator that estimates strain directly. Since the proposed estimator uses the power spectrum, it is incoherent, i.e., it does not use the phase of the signal. Previously reported incoherent methods include optical flow, speckle tracking, envelope cross-correlation, and spectral chirp z-transform techniques. Generally, incoherent methods may be less precise but significantly more robust. For example, we have demonstrated this property for the case of time-delay estimation using the envelope of echo-signals. This may be a significant advantage where elastography is to be practiced in situations involving (1) undesired scanning motion, such as the case of using an unstable handheld transducer and/or (2) undesired tissue motion, such as abdominal or intravascular elastography. This property of the estimator is demonstrated later in this paper in the simulation results section through testing of its immunity to noise caused by jitter.

[0008] The main idea behind a spectral strain approach, is based on the Fourier scaling property, which implies that a compression or expansion of the time-domain signal should lead to an expansion or compression of its power spectrum,

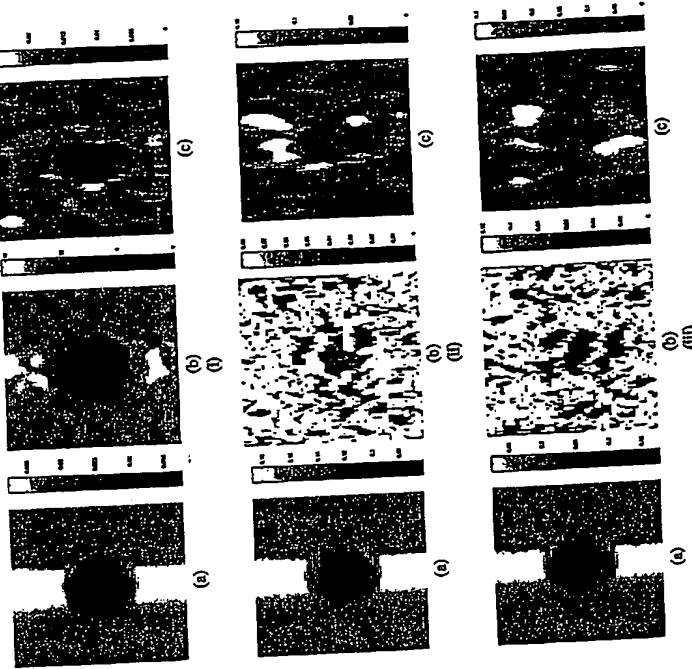


Figure 6

respectively. One of the most well known and thoroughly studied spectral motion effects is the Doppler shift, which typically links the frequency shift to the scatterer velocity between emissions. Velocities towards the transducer result in a positive frequency shift, while the opposite is true for scatterers that are move away from the transducer. However, since the scatterers within a given resolution length do not move at the same velocities, a spectrum of Doppler frequencies is observed. Therefore, initially in ultrasound, the methods of velocity estimation for the measurement of blood flow mainly operated in the frequency domain, otherwise known as spectrum analysis techniques and measured the mean velocity of scatterers across the vessel lumen (indicative of the volumetric flow rate) by estimating the mean frequency of the power spectrum. Despite the success of these techniques even in vivo vessels, detection of the Doppler frequency shift, which is typically on the order of 1 kHz, is not possible for pulsed instruments, since the downshift in frequency due to attenuation (on the order of 10–100 kHz) is expected to dominate over the Doppler shift. Since in elastography, the pre- and postcompressed segments are approximately identical depths, the attenuation effect on the two spectra is assumed to be identical and cancelled out when the two spectra are compared.

[0009] Strain estimation using spectral methods depends on the subsequent change in the scatterer statistics. Spectral methods typically link one or more signal parameters to the relative change in the mean scatterer spacing. One prior art relates the inaccuracy during a cardiac cycle. This method assumes the presence of underlying scatterer periodicities. Despite the fact that this has also been demonstrated to work in *in vivo* intravascular applications, the main assumptions of regular spacing of periodicities may not hold for most tissues. In contrast, as shown in the theory section, the spectral methods mentioned in this paper make no assumptions regarding the composition of the tissue scatterers.

[0010] Typically in elastography time-domain techniques used that involve the computation of the time-delay to estimate the displacement following an applied compression, and the estimation of strain by applying gradient operations on the previously obtained time-delay estimates. As mentioned earlier, an important advantage associated with these spectral methods as well as other estimators, such as the adaptive stretching estimator, is that they can be used to estimate strain directly; i.e., without involving the use of noisy gradient operators. In the latter case, the gradient operation introduces additional amplification of the noise into the strain estimation process, thus degrading the strain estimates. Furthermore, similar to the adaptive stretching estimator, only one estimation window is needed, for both the magnitude and the sign of the strain to be estimated.

[0011] As shown later in the theory section, spectral estimators can be divided into two main groups: a) the spectral shift methods and b) the spectral bandwidth methods. Despite the fact that we develop expressions that show direct strain estimation in both cases, in this paper we focus primarily on a spectral shift method; we estimate the relative shift in the spectral centroid caused by compressive or tensile tissue strain. Therefore, throughout this paper this new estimator is referred to as the centroid strain estimator, centroid estimator or centroid method. Current investiga-

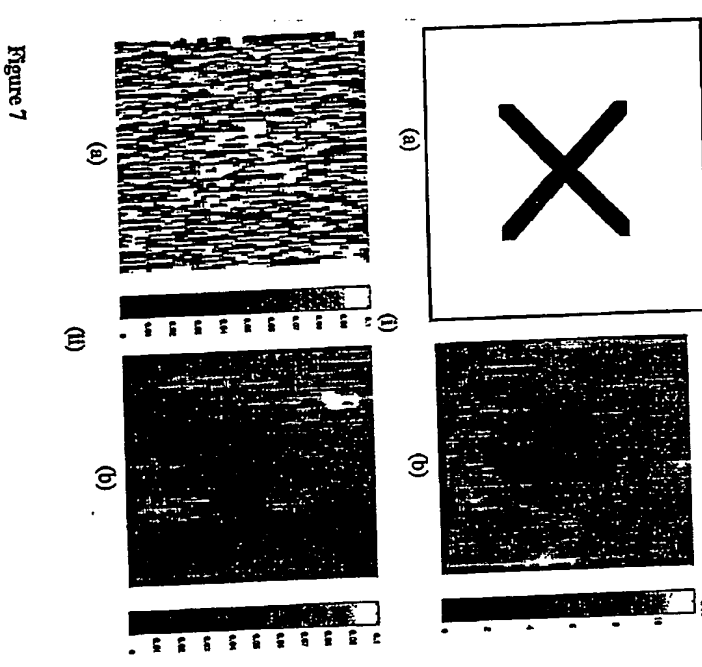
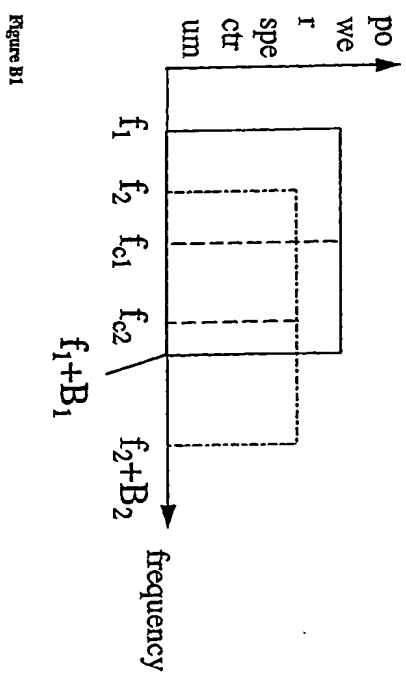
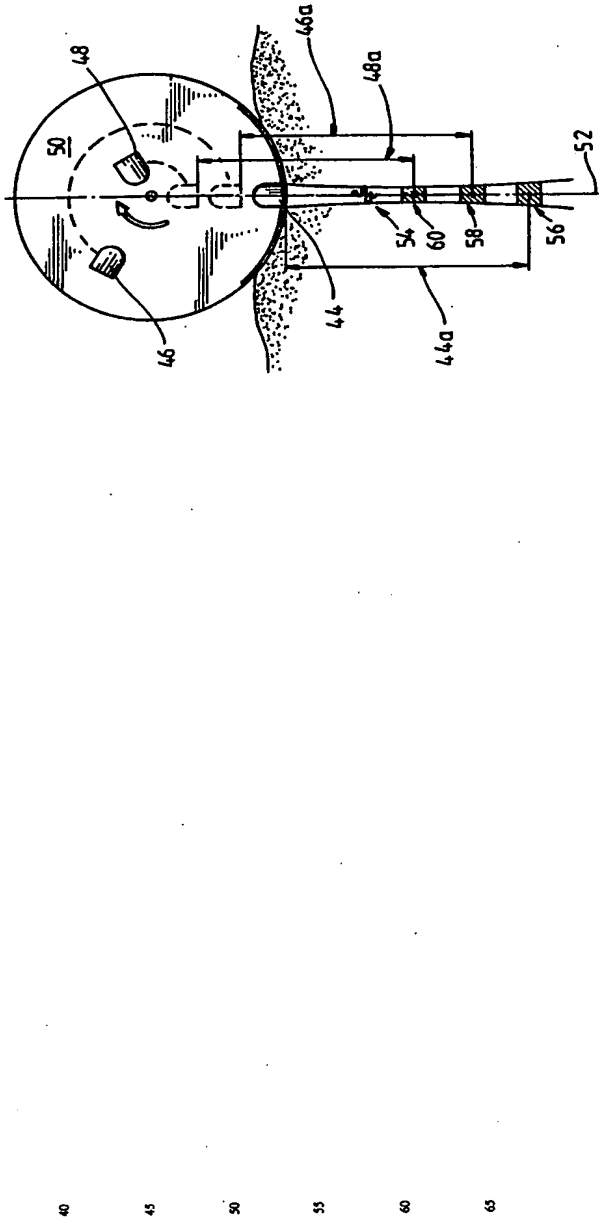
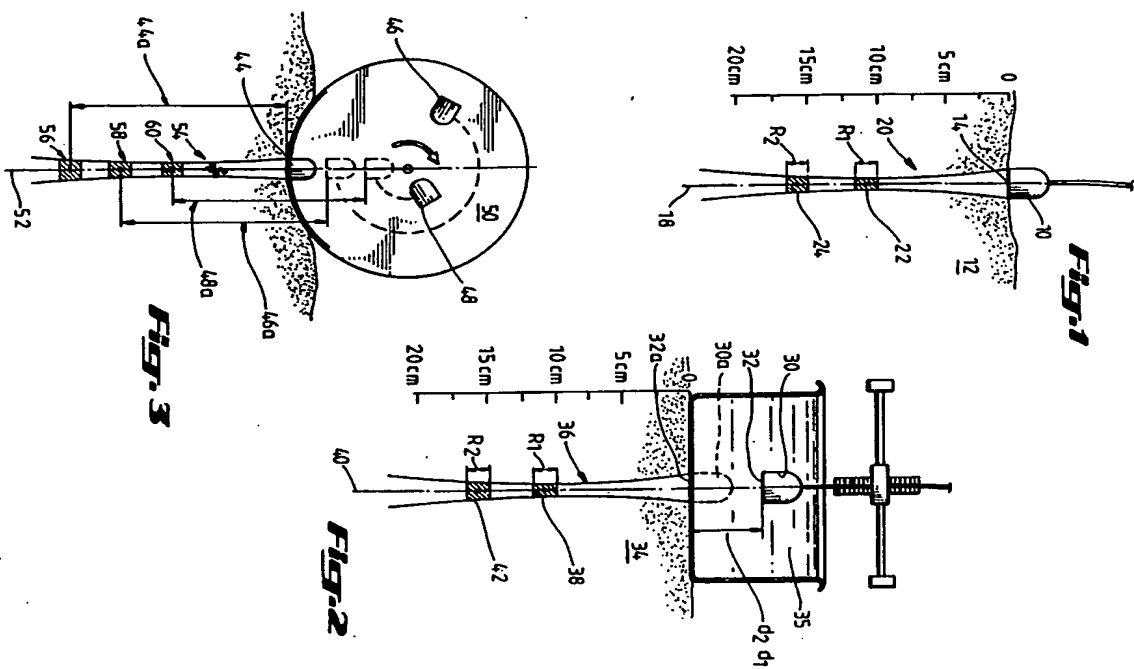


Figure 7



United States Patent [19]		Patent Number: 4,993,416	
Ophir		Date of Patent: Feb. 19, 1991	
(2) coordinating said transmissions and detections such that the signal travel times between transmission and detection for the two transducers are about equal, and (b) performing characterization measurements based upon said detected reflection signals.	13	which are axially staggered along said pulse transmission axis in a relation corresponding to the axial stagger of their respective transducers along said pulse transmission axis and (c) performing characterization measurements based upon said detected echo signals.	14
15. A method of obtaining ultrasonic echo data from target body for tissue characterization measurement which comprises:		16. The method of claim 15 comprising (a) to (d)	
(a) sequentially coupling and transmitting a separate pulse of ultrasonic energy into a target body from each one of a plurality of uncoupled ultrasonic transducers which are sequentially placed, and energized to transmit their respective pulses from, axially staggered positions along a common pulse transmission axis,		17. The method of claim 15 which further comprises performing steps (a) and (b) along a plurality of spaced transmission axes.	
(b) detecting a separate echo signal from the target body for each transmitted pulse such that the echo signals emanate from regions within the tissue		18. The method of claim 11, wherein said characterization measurement performing step comprises obtaining the attenuation coefficient of the target body by the log spectral difference method.	
19. The method of claim 18 in which adjacent said axes are spaced sufficiently from each other to be non-correlatable.		• • • •	
OTHER PUBLICATIONS		Elimination of Diffraction Error in Acoustic Attenuation Estimation Via Axial Beam Translation, Ultrasonic Imaging, vol. 10, pp. 139-152 (1988), by Ophir and Melhaft.	
Primary Examiner—Frances Jaworski Attorneys, Agents, or Firm—Arnold, White & Durkee		[57] ABSTRACT	
[54] SYSTEM FOR ULTRASONIC PAN FOCAL IMAGING AND AXIAL BEAM TRANSLATION	[75] Inventor: Jonathan Ophir, Houston, Tex.	The present invention provides a novel method and apparatus which allows concurrent imaging and rapid axial beam translation measurements used to calculate the attenuation characteristics of a target body. The present invention employs an ultrasonic scanner which contains a plurality of matched transducer elements. These elements are staggered on a meander path which sequentially places each transducer opposite an acoustic window at axially spaced positions along a common axis. The present invention also enables axial beam translation techniques to be adapted to current ultrasonic imaging systems.	
[73] Assignee: Regents The University of Texas System, Austin, Tex.	[21] Appl. No.: 343,405	[51] Int. Cl. G01K 1/00	
[22] Filed: Apr. 25, 1989	[52] U.S. Cl.	[58] Field of Search ... 128/660.06, 660.09	
		128/660.1; 73/631, 639, 599	
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19 Claims, 4 Drawing Sheets		[51] U.S. CL. 128/660	





7. Optimize the data from step 6 to obtain the frequency band width that provides the best linear regression with frequency using a standard deviation parameter; and

8. Compute the attenuation coefficient.

Although the invention has been described with a certain degree of particularity, it is to be understood that the above description has been only by way of example. Numerous other changes will be apparent to those reading the specification without departing from the spirit and scope of the invention as claimed.

What is claimed is:

1. Apparatus for tissue characterization measurement of a target body comprising:

(a) a plurality of matched ultrasonic transducers; (b) a moveable mounting member adapted to mount the transducers in a spaced array such that movement of the mounting member sequentially positions the transducers in an axially staggered pattern along one or more common ultrasonic radiation axes to transmit and receive ultrasonic signals to and from the target body along the one or more common ultrasonic radiation axes up, and

(c) means coupled to said matched transducers for performing tissue characterization measurements using said received ultrasonic signals.

2. The apparatus of claim 1 wherein said mounting member sequentially positions the transducers along a plurality of common ultrasonic radiation axes.

3. The apparatus of claim 2 comprising four to six matched transducers.

4. Apparatus for ultrasonic tissue characterization measurement of a target body comprising:

(a) a plurality of matched ultrasonic transducers, each of said transducers having an ultrasonic aperture; (b) a moveable mounting member adapted to mount the transducers in a spaced array such that movement of the mounting member sequentially positions the transducers in an axially staggered pattern along one or more common ultrasonic radiation axes to transmit ultrasonic signals to the target body along the one or more common ultrasonic radiation axes;

(c) a range gate operable to detect a separate ultrasonic echo signal originating from said target body; (d) in response to each transmitted signal wherein: (1) the detected echo signals emanate from regions within the target body which are axially staggered along said radiation axes in a relation corresponding to the axial stagger pattern of the transducers; (2) along said axes, and (2) the distances between each of said regions and the ultrasonic spotters of their respective transducers are about equal.

(e) means coupled to said matched transducers for performing tissue characterization measurements using said received ultrasonic signals.

5. The apparatus of claim 2 wherein said mounting member sequentially positions the transducers along a plurality of common ultrasonic radiation axes.

6. The apparatus of claim 5 comprising four to six matched transducers.

7. Apparatus for ultrasonic characterization measurement of a target body comprising:

- (a) a sealed housing adapted to contain an acoustic coupling fluid and having an acoustically transparent window disposed on the periphery;
- (b) a disk disposed inside said housing and rotatably coupled to the housing;

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11. The apparatus of claim 7 comprising:

(a) a plurality of matched transducers mounted to face radially outward on the disk in a spirally staggered pattern such that rotation of the disk sequentially and sequentially places said transducers across said window at axially spaced positions along one or more common ultrasonic radiation axes; and

(b) means coupled to said matched transducers for performing tissue characterization measurement.

8. The apparatus of claim 7 comprising:

9. The apparatus of claim 8 wherein said transducers are focused transducers.

10. The apparatus of claim 5 wherein said mounting member sequentially positions the transducers along a plurality of common ultrasonic radiation axes.

11. An apparatus for ultrasonic analysis of a target body comprising:

(a) a housing adapted to make contact with said body and to contain an acoustic coupling fluid;

(b) a mounting member mounted within the housing in rotatable relation thereto such that rotation of the mounting member causes points along the periphery of the mounting member to travel past the portion of the housing which makes contact with said target body;

(c) a plurality of matched ultrasonic transducers mounted on the mounting member in a pattern such that movement of the mounting member sequentially positions the transducers each transducer adjacent the sequentially positioned transducers and spaced along a common ultrasonic radiation axis extending through the contact into the body; and

(d) circuitry operable to separately activate each transducer when adjacent said contact so as to (1) transmit ultrasonic energy into the body and therefrom; (2) receive a reflection of said energy from a region within the body such that the travel times of the energy between the transducers and their respective reflective regions within the body are about equal.

12. The apparatus of claim 11 wherein the transducers are mounted in a spiral pattern on the moveable member.

13. The apparatus of claim 6 which further comprises an acoustic window located in the portion of the housing which is adapted to contact the body.

14. A method of performing an ultrasonic characterization measurement of a target body using a plurality of ultrasonic transducers which comprise:

(a) sonically coupling a first ultrasonic transducer to a target body to transmit and receive ultrasonic signals to and from the body along a radiation axis;

(b) transmitting an ultrasonic signal from the first transducer along said radiation axis into the body;

(c) detecting reflection signals of the transmitted signal reflected from the body during a selected time interval following the transmission of the transmitted signal;

(d) repeating steps (a) through (d) for the second transducer;

(e) moving and moving the first ultrasonic trans-

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12. The apparatus of claim 11 which further comprises:

(a) a plurality of matched transducers mounted to face radially outward on the disk in a spirally staggered pattern such that rotation of the disk sequentially and sequentially places said transducers across said window at axially spaced positions along one or more common ultrasonic radiation axes and

(b) means coupled to said matched transducers for performing tissue characterization measurement.

9. The apparatus of claim 8 wherein said transducers are focused transducers.

10. The apparatus of claim 5 wherein said mounting member sequentially positions the transducers along a plurality of common ultrasonic radiation axes.

11. An apparatus for ultrasonic analysis of a target body comprising:

(a) a housing adapted to make contact with said body and to contain an acoustic coupling fluid;

(b) a mounting member mounted within the housing in rotatable relation thereto such that rotation of the mounting member causes points along the periphery of the mounting member to travel past the portion of the housing which makes contact with said target body;

(c) a plurality of matched ultrasonic transducers mounted on the mounting member in a pattern such that movement of the mounting member sequentially positions the transducers each transducer adjacent the sequentially positioned transducers and spaced along a common ultrasonic radiation axis extending through the contact into the body; and

(d) circuitry operable to separately activate each transducer when adjacent said contact so as to (1) transmit ultrasonic energy into the body and therefrom; (2) receive a reflection of said energy from a region within the body such that the travel times of the energy between the transducers and their respective reflective regions within the body are about equal.

12. The apparatus of claim 11 wherein the transducers are mounted in a spiral pattern on the moveable member.

13. The apparatus of claim 6 which further comprises an acoustic window located in the portion of the housing which is adapted to contact the body.

14. A method of performing an ultrasonic characterization measurement of a target body using a plurality of ultrasonic transducers which comprise:

(a) sonically coupling a first ultrasonic transducer to a target body to transmit and receive ultrasonic signals to and from the body along a radiation axis;

(b) transmitting an ultrasonic signal from the first transducer along said radiation axis into the body;

(c) detecting reflection signals of the transmitted signal reflected from the body during a selected time interval following the transmission of the transmitted signal;

(d) repeating steps (a) through (d) for the second transducer;

(e) moving and moving the first ultrasonic trans-

9 **9** gated strips 76 & seq. consists of a series of range gated regions such as 92. The number and location of these regions corresponds to the number of axes along which a transducer is pulsed as it sweeps the acoustic window. The number of times a transducer is pulsed corresponds to the radial scan line or a-line density. An a-line is the echo sequence corresponding to an ultrasound pulse as it propagates along a fixed axis through the target 95. The desired a-line density is based on considerations of adequate imaging and depth of penetration, and may be, for example, 170 a-lines per  $\pi/3$  sector angle 96, for a depth of penetration of 20 centimeters. These figures, combined with the average speed of sound in tissue of 1540 meters per second, result in a sweep time per  $\pi/3$  sector angle 96 of 45 milliseconds, or about 0.27 seconds 15 for one revolution of disk 62. This allows a frame rate of approximately four frames per second. This allows combined with hand-held contact scanning allows convenient imaging and attenuation estimation with A/B/C. Continuing in FIG. 5, as transducer 76 sweeps across acoustic window 66, it is activated approximately 170 times along 170 axes and acquires echoes corresponding to 170 regions (temporal windows) to form strip 76a. For example, an acoustic pulse will propagate within a parabolic beam having an axis 96. An echo signal which corresponds to gated strip 92 is acquired from the a-line propagating along axis 96 using a range gate. As each successive transducer sweeps the acoustic window, each will be pulsed along the same axes, including 96, 30 and acquire echo signals from its respective range gated strip, including 96, for ex. seq. The data from the acquired signals may be combined to generate an extended focus image or may be used for attenuation estimation. Because of the transducer staggered pattern, each range gated strip is at a constant distance from its respective transducer aperture.

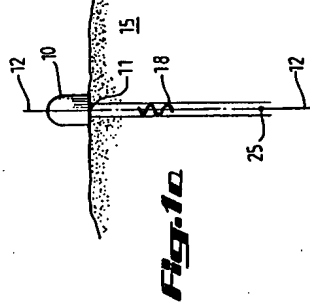
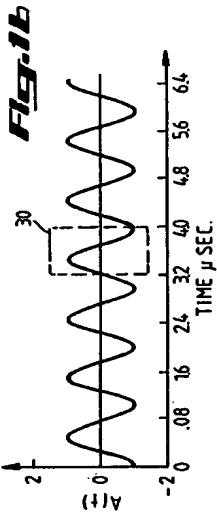
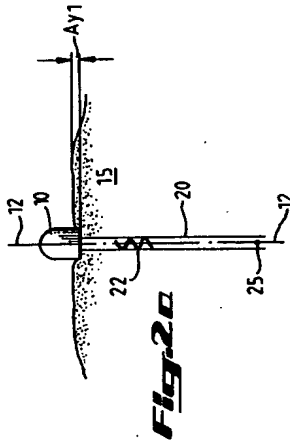
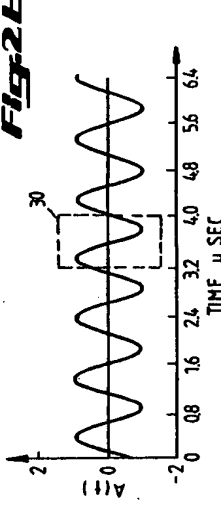
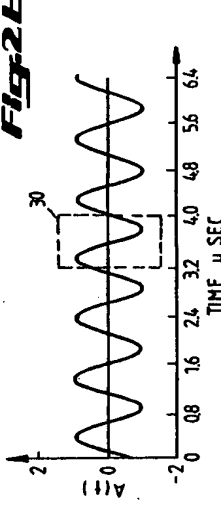
By requiring approximately 170 echo sequences (a-line) from the extended focus sector 96, the apparatus of FIG. 5 may obtain good quality images. However, in order to obtain a good measurement of the attenuation characteristics, the echo sequences used are advantageously "uncorrelated". A "correlated" a-line represents a non-statistically independent echo sequence. For example, at a range of 100 millimeters, 170 echo sequences across the  $\pi/3$  sector angle correspond roughly to two a-lines per millimeter, and thus would be considered highly "correlated". This means that about only 1 out of 10 echo sequences should typically be used for attenuation estimation, or about 17 echo sequences per 30 frame.

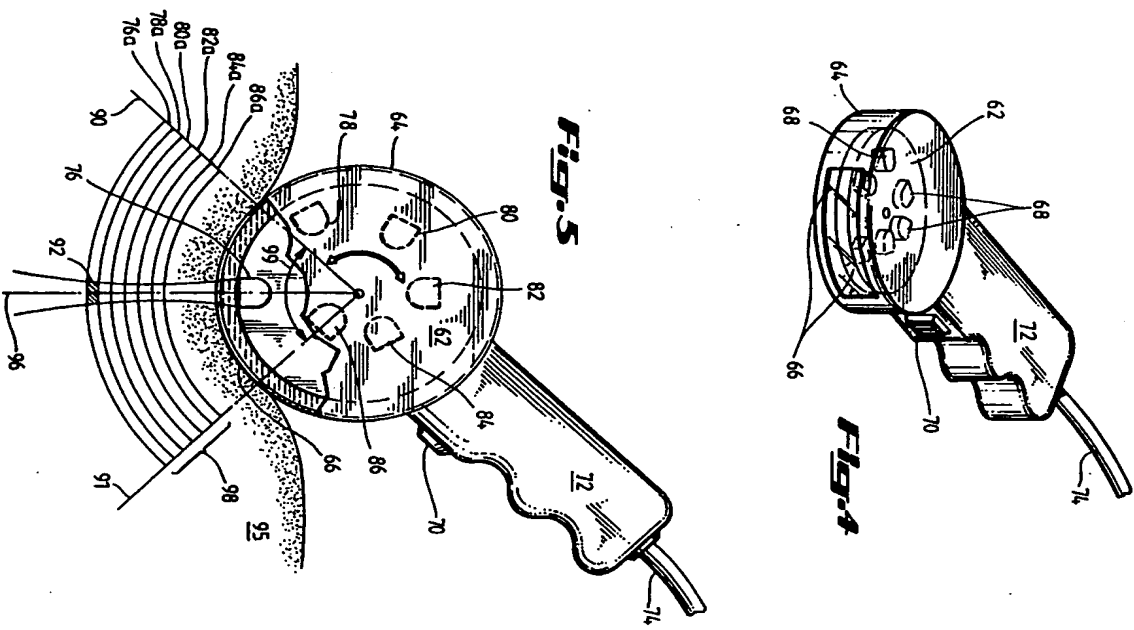
Typically, about 20 independent echo sequences may be acquired for an attenuation estimation. Conveniently, the a-lines may be acquired from multiple uncorrelated planes or frames. Therefore, the use of the scanner head *in vivo* may involve rocking the scanner from side to side over an angle of  $\pi/4$  or so, and acquiring data from about 10 "uncorrelated" frames. If the device generates approximately 4 frames per second, rocking of the transducer will be accomplished in about 2.5 seconds. During the first frame, all 170 echo sequences (each comprising 6 segments acquired by the axially staggered transducers) may be digitized at a 25 MHz sampling rate and stored in half of the data acquisition system 116 in FIG. 6. During the second frame the echo data sequences may be stored in the other half of the data acquisition system 116, while the data from the first frame is transferred to the computer 130.

10 Referring to FIG. 6, a representative block diagram of a system implementing the present invention is shown. The A/B/C scanning head 100 is driven by a 1024 step/revolution stepper motor 106, whose speed is controlled by a precision 3.9 KHz pulse repetition frequency ("PRF") clock 102, derived from a higher frequency 2 MHz master clock 104. An optical shaft encoder 108 is connected to the stepper motor 106 and provides absolute shaft angle information with 11 bits of accuracy. The three most significant bits may serve as sector identifiers. The remaining bits may be used to code the individual echo sequences or a-lines. As the scanning head turns, the transducer multiplexer 110 sequentially selects one of the six transducers which is to be driven by the transmitter 112. Both the transmitter and the time-gain-compensation ("TGC") circuit 114 are driven from the 3.9 KHz PRF clock 102. The TGC or depth-compensation circuit amplifies echo signals in proportion to their transit time to compensate for signal attenuation. This allows for improved images on the display 118 and complete digitization of signals when fed into the data acquisition and storage system 116. A range gate 120 provides a variable signal to the digital scan converter ("DSC") 122 which allows pixels to be modulated on the display 118 corresponding to their respective range gated strips. The signal intended for imaging is fed into the demodulator 124 and then fed to the DSC 122. Position signals to the DSC 122 are provided by a dual XY programmable-read-only-memory ("PROM") 126 which is strobed by the 2 MHz clock 104. The DSC 122 generates a sequence of pre-programmed XY addresses at a rate of 2 MHz. These addresses are converted to analog (position) signals and fed to the DSC 122. The DSC 122 may operate in gated survey mode, such that individual sequential range gated strips in the image are updated, corresponding to the transducer with the appropriate focal distance which is swept across the acoustic window.

The A/B/C section of the exemplified apparatus involves eight bit digitization of the radio frequency ("RF") signal at 25-50 MHz via a data acquisition and storage system 116, and the TGC voltage at 25-50 KHz. These signals are later combined in software to calculate the absolute magnitude of the echo spectra. The output of the analog-digital converter 128 is communicated to the microcomputer 130 through the IEEE-488 bus 132. Using a fast transducer scanner, the software for computing the attenuation coefficient may comprise, for example:

1. Acquire 170 a-lines and divide into 6 segments (range gated strips) corresponding to 6 depths of observation within the target body;
2. Compute the average power spectrum for each depth by averaging the squared fast Fourier transform for all echoes in each segment over all a-lines;
3. Perform spectral smoothing by windowing the autocorrelation function of the averaged spectra obtained in step 2 above and re Fourier transforming to obtain a smoothed power spectrum;
4. Convert the power spectra of step 3 to units of dB (log of power spectrum);
5. Perform linear regression with respect to each depth of observation for all frequencies in the ultrasonic pulse, (wideband) to obtain an attenuation-with-frequency curve;
6. Determine the best frequency band within the bandwidth of frequencies in step 5 to obtain a linear fit with respect to frequency;

**FIG. 10****FIG. 11a****FIG. 11b****FIG. 2a****FIG. 2b**



**7** radiation axis 52. As the scanner mechanism turns, transducer 46 is aligned at a separate point along axis and acquires echoes from region 60. Similarly, transducer 46 is positioned at a unique point along axis 52 and acquires echoes from region 58. As will be appreciated from FIG. 3, the distances 46a, 46c, and 46e between the apertures of transducers 46, 46 and 46 and their respective regions are identical. In this way, the elements of ABT are achieved, but without the necessity of a bulky water bag. By using matched transducer, <sup>10</sup> f--i.e., transducers having substantially identical acoustic properties, the scanner of the present invention achieves the same results as a single transducer axially translated along a common axis in a water bag as in FIG. 2. It will also be appreciated that the speed and timing of slow data acquisition which is undesirable in clinical settings.

The present invention may advantageously perform ABT techniques with a scanner as shown in FIG. 4. The scanner comprises a disk 62 rotatably disposed in a sealed housing 64 which contains an acoustic window 66. The acoustic window is sonot-permeable and may be constructed from a membrane of a high strength thermoplastic resin, such as a polycarbonate or the like. The housing is filled with an acoustic coupling fluid which is matched to the speed of sound and impedance of the target body. The disk 62 contains a plurality of spatially staggered and matched transducer elements 68. As the disk rotates, different transducer will scan the acoustic window 66 at slightly different positions along one or more common axes. Because of the staggering of the transducers 68, sequential transducer sweeps will be operable to acquire echo signals from sequentially staggered regions within a target body. As will be apparent from FIG. 4 and FIG. 5, the regions may be staggered both axially and circumferentially.

ABT may be achieved by appropriate range gating of the returned echoes such that only echoes returning from regions at a fixed distance from each respective 20 transducer element are received. The range gate may be set, for example, to acquire echo signals corresponding to regions having a width of about 1 cm to about 3 cm. These echoes may be used to generate an image or may be used to calculate the attenuation coefficient. 25

The diameter of the scanner is usually somewhat larger than twice the depth of the body region of interest targeted for attenuation estimations. In clinical diagnosis, the body contact area may range from about 6 cm  $\times$  2 cm to about 9 cm  $\times$  2 cm, depending on the number of spatially staggered transducers employed. The apparatus may optionally contain a manual ABT switch 70 located in handle 76 of the scanning mechanism. This allows the operator to obtain an automatic range

before activating the ABT mode. Electrical lead 76 may connect the scanner of FIG. 4 to an ultrasonic diagnosis system such as exemplified in FIG. 6.

The apparatus and method of the present invention require a plurality of spatially staggered and matched transducers to achieve the advantages of quick, acoustic rate and non-problematic attenuation measurements. Optionally, additional unsteered transducer elements may be included for conventional imaging only. The invention contemplates transducers which may be piezoelectric, ferroelectric or magnetostriuctive in nature.

The present invention is not limited by the size, focusing properties or band width of the transducer elements to be employed.

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### 8

In general a focused transducer has an ultrasonic beam which in a certain range is constricted or narrow in diameter than the fixed diameter beam generated by a non-focused transducer. The range from the transducer aperture at which a focused beam is constricted is known as the working range of the transducer. Outside of this range the focused beam is more divergent than an unfocused beam. In one embodiment of the present invention, a plurality of focused transducers may be used. Sequential transducer sweeps across the acoustic window then have sequentially staggered focal lengths. By accepting signals only from the focal (working) range of each transducer element and combining the images so obtained, ABT may be achieved and high resolution imaging in the extended focus may also be simultaneously done. However, the present invention contemplates the use of either focused or unfocused transducers.

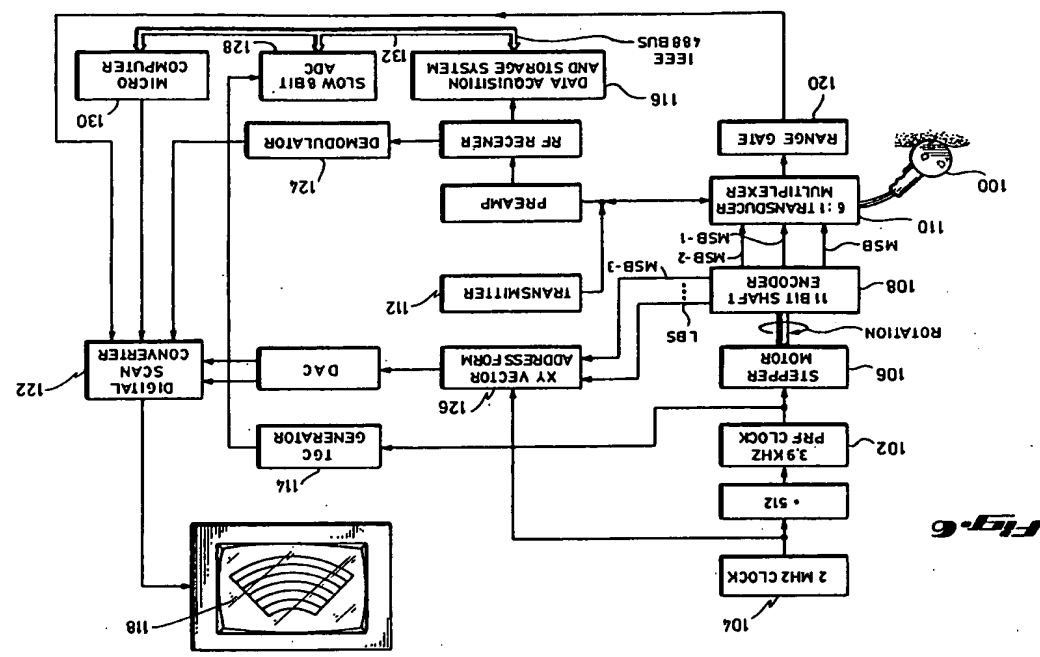
The apparatus and method of the present invention are not limited to a particular algorithm for calculating the attenuation characteristics of a target body. The present invention optionally contemplates using a combination of ART and IDP algorithms to further correct echo spectra obtained during attenuation measurement. For a review of IDP algorithms and techniques, see Cordonio, et al., "Diffraction Correction in Pulse Echo Attenuation Measurement", IEEE Transac. Symp. Proc. 841-846 (IEEE Cat. No. 83 Ch. 194-1, 1983); Cloostermans, et al., "A Beam-Corrected Estimation of the Frequency Dependent Attenuation of Biological Tissue from Backscattered Ultrasound", Ultrasonic Imaging, Vol. 1, 136-147 (1982); and Imano, et al., "Improvements in the Spectral Difference Method for Measuring Ultrasonic Attenuation", Ultrasonic Imaging, Vol. 5, 331-345 (1983), which are incorporated by reference herein.

For a further review of algorithm used in attenuation characterization, see Leeman, et al., "Perspectives on Attenuation Estimation from Pulse-Echo Signals," IEEE Transactions on Sonics and Ultrasonics, Vol. SU-31, No. 4, 352-361 (1984) and Gara, et al., "In Vivo Attenuation Measurement Method and Clinical Reference," Proc. 6th European Communities Workshop on Acoustic Imaging, Vol. 1, 136-147 (1982); and Imano, et al., "Improvements in the Spectral Difference Method for Measuring Ultrasonic Attenuation", Ultrasonic Imaging, Vol. 5, 331-345 (1983), which are incorporated by reference herein.

Although the apparatus and method of this invention are typically described in relation to clinical diagnosis, this should be understood not to be a limiting factor on the utility of the invention. To the contrary, the present invention has utility in any area in which the attenuation characteristics of a target body may be desired. For example, the present invention may be used in forensics, tissue characterization studies, veterinary medicine, laboratory experiments or measuring the properties of any material which exhibits attenuation and scattering of ultrasonic energy.

As schematically shown in FIGS. 4 and 5, an ultrasonic scanner having six spatially staggered transducer elements 76, 78, 80, 82, 84 and 86 which correspond to a target body 98. The extended focus 96 of the body 98 having boundaries 90 and 91 is diagnosed as the transducers sequentially sweep acoustic window 66. The extended focus sector 98 comprises six range gated transducers, having six spatially staggered transducer elements 76, 78, 80, 82, 84 and 86 which correspond to a target body 98. The extended focus 96 of the body 98 having boundaries 90 and 91 is diagnosed as the transducers sequentially sweep acoustic window 66.

The extended focus sector 98 comprises six range gated transducers, having six spatially staggered transducer elements 76, 78, 80, 82, 84 and 86 which correspond to a target body 98. The extended focus 96 of the body 98 having boundaries 90 and 91 is diagnosed as the transducers sequentially sweep acoustic window 66.



In (D2)  $d = R - W$  is the portion of  $R$  for which  $\alpha$  is non-zero, and  $W$  is the portion for which  $\alpha$  is zero. Assuming that there are two scattering ensembles at ranges  $R_1$  and  $R_2$ , respectively, having the same average value for  $\langle \tau^2 \rangle$  and both within the attenuating target body (see FIG. 4, regions 22 and 24), then the ratio of the received ultrasound echo from both ranges is:

$$(D3) \quad P_r(R) = \exp\left(\frac{4\pi(d_1 - d_2)}{R^2}\right)$$

The variables  $d_1$  and  $d_2$  may be expressed as  $R - W$  and  $R_2 - W$  respectively. The constant speed of sound in the water path mechanism is assumed to be the same as that of the target body tissue. The wavelength of the ultrasound pulse is also assumed constant. The numerator of (D3) contains the desired attenuation information, whereas the denominator is a beam-spreading loss which causes artifacts in acquired echoes and results in bias errors in estimating the attenuation characteristics recorded as a function of frequency to derive the attenuation coefficient. However, the attenuation coefficient which is thereby derived, will normally be subject to bias errors. Artifacts in the spectra calculated from echoes originating from regions at different distances from the transducer aperture 14 often result from beam-spreading loss, diffraction and/or interference effects. These variations in transducer pulse-echo impulse response as a function of distance from the aperture contaminate the spectra. The data from these spectra will subject attenuation estimations thereby derived to bias errors.

FIG. 2 exemplifies the use of AFT to reduce the bias errors in the attenuation estimation discussed above.

Transducer 30 is shown disposed in water path 35 which is optically coupled to body 34. Echoes are acquired from region 38 in response to acoustic beam 36 using range gating techniques as set forth above. The transducer 30 is then translated along the radiation axis 40 to position 38 wherein echoes from region 42 are acquired. The distance between aperture 32 and region 38 is identical to the distance between the aperture at 32 and region 42. By keeping the distance between each transducer aperture and its respective region of interest constant during echo acquisition, the bias errors in the derived attenuation coefficient may be reduced.

The present invention utilizes AFT techniques to reduce bias errors in attenuation estimations without the need for a bulky water bag. According to the present invention, AFT may be conveniently performed by utilizing a plurality of matched transducers mounted on a scanning mechanism in an axial stagger pattern. As a result of the pattern, the mechanism will sequentially place each respective transducer across an acoustic window at axially different positions along one or more common axes.

Conveniently, the present invention may employ a rotating scanning head which carries a plurality of transducers A as illustrated in FIG. 3. transducer 44, 46 are mounted on a rotor 50 and face radially outward in a spirally staggered pattern. As the rotor 50 turns, the transducers are sequentially positioned at different points along the axis 52 and thus become axially staggered. As each transducer sweeps past the axis 52, an ultrasonic pulse 54 is generated and echo spectra acquired for temporal windows corresponding to regions 56, 58 and 60. For example, transducer 44 is used to acquire echoes from region 56 along the ultrasonic beam

toward the aperture 14 of transducer 10. Using known range-gating techniques, the ultrasonic echo sequence is broken up into temporal windows which may correspond, for example, to regions 22 and 24 within body 12. Since ultrasound propagates through soft tissue at about 1500 meters per second, it will take ultrasound about 1.30 microseconds to go and return through 10 cm of tissue. Thus, a range gate set to receive echoes for a 130-156 microseconds temporal window will acquire 10 echoes corresponding to region 22. A range gate set for 195-221 microseconds will acquire echoes corresponding to region 24. The echoes are converted to a series of spectra using known Fourier transform algorithms. See Roman Kuc, "Estimating Acoustic Attenuation from Reflected Ultrasonic Signals: Comparison of Spectral Shift and Spectral Difference Approaches," IEEE Transactions on Acoustics, Speech, and Signal Processing, ASSP-32, 1-6 (1984). The log-spectral differences between each region may be computed and plotted or recorded as a function of frequency to derive the attenuation coefficient. However, the attenuation coefficient which is thereby derived, will normally be subject to bias errors. Artifacts in the spectra calculated from echoes originating from regions at different distances from the transducer aperture 14 often result from beam-spreading loss, diffraction and/or interference effects. These variations in transducer pulse-echo impulse response as a function of distance from the aperture contaminate the spectra. The data from these spectra will subject attenuation estimations thereby derived to bias errors.

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Roman Kuc, "Estimating Acoustic Attenuation from

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## SYSTEM FOR ULTRASONIC PAN FOCAL IMAGING AND AXIAL BEAM TRANSLATION

### BACKGROUND OF THE INVENTION

#### 1. Field of the Invention

This invention relates generally to methods and apparatus for performing ultrasonic diagnosis of a target body and more particularly to methods and apparatus for concomitantly acquiring an ultrasonic image and measuring the attenuation characteristics of a target body.

#### 2. Description of Related Art

Traditional ultrasonic diagnosis is performed by transmitting ultrasonic energy into a target body and generating an image from the resulting echo signals in order to survey anatomical structures. A transducer is used to both transmit the ultrasonic energy and to receive the echo signals. During transmission, the transducer converts electrical energy into mechanical vibrations. Acquired echo signals produce mechanical oscillations in the transducer which are converted to electrical signals for amplification and recognition.

A human or animal body represents a non-homogeneous medium for the propagation of ultrasound energy. Images may be generated from the echo backscatter signals which are produced as ultrasonic energy propagates through acoustic impedance interfaces and scatter sites within the target body. A scatter site within the target area is measured from the acoustic impedance changes at boundaries of varying density and/or sound speed within a target body. A knowledge of the speed of sound in tissue permits the determination of the depth and location of both the interfaces and the scatter sites from the measurement of echo travel time. An image may be generated from this information by modulating the intensity of pixels on a display in proportion to the intensity of echoes from corresponding points in the target.

In recent years, much effort has been expended to obtain clinically useful data from the several physical techniques involved in echo production. In particular, techniques for measuring the attenuation of ultrasonic energy as it propagates through a soft tissue have been pursued. An ultrasonic energy propagates through tissue, some of the energy is absorbed and some is scattered out of the acoustic pathway. These two mechanisms result in a net loss of signal power—i.e., acoustic attenuation. Acoustic attenuation has a near linear frequency dependence and is generally characterized as a power law in decibels per centimeter of propagation per MHz. When attenuation is plotted as a function of frequency, a near linear relationship is defined. This function—the attenuation coefficient—varies with the acoustic characteristics of particular tissues. In soft tissue it has been shown that there is some correlation between attenuation characteristics and tissue pathlength. For example, a healthy liver will have a different attenuation coefficient than a cirrhotic liver. Thus, a technique for accurate measurement of acoustic attenuation characteristics would be clinically valuable.

Several prior art techniques have been developed to measure the attenuation characteristics of soft tissue. One prior art technique for measuring ultrasound attenuation requires that the target body be placed between separate transmitting and receiving transducers. This technique, however, is impractical in clinical settings, since transmission through most sections of a target body is not possible. A second prior art technique known as the "substitution method" utilizes a known reflector to return echoes to a common transmitting/receiving transducer. A tissue specimen is placed between the transducer and the reflector, and the resulting decrease in echo signal power is used to determine the attenuation characteristics of the tissue. This technique, however, also has clinical limitations, since known reflectors cannot be inserted into human target bodies.

Both of the above prior art techniques are limited by the measurement of attenuation based upon the entire path of the propagating energy. As such they are not entirely suitable for measuring attenuation of limited regions of interest within a target body.

Knowledge of the ultrasound attenuation characteristics within limited regions of a human or other animal target body has specific value for clinical diagnostic purposes. As discussed above, for example, diseased liver have different attenuation characteristics from healthy one. A technique known as the "spectral difference" method has been proposed, whereby echo signals from varying depths in the target are acquired and converted to spectral signals using known Fourier transform algorithms. See Roman Kuc, "Estimating Acoustic Attenuation from Reduced Ultrasonic Signals: Comparison of Spectral Shift and Spectral Difference Approaches", IEEE Transactions on Acoustics, Speech and Signal Processing, ASSP-32, 1-6, (1984). A knowledge of the speed of sound in the target allows the ultrasonic echo sequence to be broken up into temporal spectral windows which correspond to varying depths in the target. The log-spectral differences between windows down are computed, and the attenuation coefficient is derived. However, attenuation estimations using this technique tend to suffer from bias errors. These errors are attributable to a variety of phenomena, including the inherent differences in pulse-echo impulse response of a transducer at various distances from the transducer.

One method for eliminating the effects of these beam diffraction errors has used Inverse Diffraction Filtering ("IDF") techniques to develop a beam correction factor in clinical settings since pulse-echo beam char-

Imaging '70, 139-152 (1968) which is incorporated by reference herein. However, initial ABT techniques have tended to be problematic in clinical settings due to the requirement for axial translation of the transducer in a bulky water bag. Also, these initial ABT techniques require several minutes to collect sufficient ultrasonic echo data for accurate attenuation estimations. This time requirement is not very desirable for current medical diagnosis purposes. Thus, until the present invention, ABT techniques have not been adaptable to current ultrasonic imaging systems.

Several forms of ultrasonic scanning mechanisms have been suggested for use in performing ultrasonic diagnosis of human and animal organs. One principal form employs a rotating scanning head which carries a plurality of transducers. As the head rotates, the transducers sequentially pass by a body organ at which time they transmit signals into the organ and receive reflected echoes from the organ.

A second principal form of ultrasonic scanning mechanism employs an oscillating scanning head which carries a single transducer. As the transducer oscillates through a scanned angle while transmitting and receiving ultrasonic signals.

In both the rotating and oscillating scanning mechanism, the scanning head may be mounted in a housing filled with an acoustic coupling liquid. In some instances, the motor driving the head may be mounted directly in the same liquid-filled housing; in other instances, it may be mounted in a dry region outside the housing and coupled to the head through a suitable seal.

### SUMMARY OF THE INVENTION

The method and apparatus of the present invention address the problems inherent in the prior art, including the bulky water bag and the slow data acquisition of the known ABT methods. The present invention provides a novel method and apparatus which allows concurrent imaging and rapid ABT measurements. The present invention also enables ABT to be adapted to current ultrasonic imaging systems.

The present invention comprises a method and apparatus for concomitant ultrasonic pan focal imaging and axial beam translation. The present invention employs an ultrasonic scanner which contains a plurality of matched transducer elements. These elements are staggered on a mechanism which sequentially places each transducer opposite an acoustic window at stably spaced positions along a common ultrasonic radiation axis. Further, the mechanism may scan across the window so that the transducers may repeat this axial orientation along a plurality of such radiation axes which radiate through the same window. Thus, the invention comprises a system for scanning a region within a tissue of unknown tissue. See Robinson, et al., "Beam Pattern (Diffraction) Correction for Ultrasonic Attenuation Measurements", Ultrasonic Imaging, Vol. 6, No. 3, 291-303 (1984). Thus, the IDF correction factor to be applied to echo signals varies according to the type of (generally unknown) tissue being examined.

A second method for elimination of diffraction errors known as Axial Beam Translation ("ABT") has been successful in rendering unbiased estimations of attenuation in unknown targets. See Ophir and Mehta, "Elimination of Diffraction Error in Acoustic Attenuation Estimation Via Axial Beam Translation", Ultrasonic

Because of the axial staggering of the elements, sequential transducer scans may acquire echo data emanating from successively deeper or shallower regions in the target body corresponding to the stagger pattern.

FIG. 2 is a schematic view useful for explaining the effective aperture of the transducer at frequency  $\Omega = c/\lambda$ .

FIG. 3 is a schematic view useful for explaining an understanding of the invention may be obtained by referring to the following detailed description when taken in conjunction with the accompanying drawings.

FIG. 4 is a perspective interior view of an ultrasonic scanner assembly useful for explaining an understanding of the invention when taken in conjunction with the following detailed description when taken in conjunction with the accompanying drawings.

FIG. 5 is a schematic anterior view of the ultrasonic scanner assembly of FIG. 4 shown acoustically coupled to a target body.

FIG. 6 is a block diagram of an ultrasonic diagnostic system according to one embodiment of the present invention.

### DETAILED DESCRIPTION

The Axial Beam Translation ("ABT") method relies on the acquisition of echoes from varying depths in the target body, while maintaining a constant range from the transducer aperture. The ratio between the received (echo) and transmitted (pulse) of ultrasonic power for matched transducer elements. These elements are staggered on a mechanism which sequentially places each transducer opposite an acoustic window at stably spaced positions along a common ultrasonic radiation axis. Further, the mechanism may scan across the window so that the transducers may repeat this axial orientation along a plurality of such radiation axes which radiate through the same window. Thus, the invention

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$$\frac{P_{ABT}}{P_0} = T^2 \cdot e^{-2d} \cdot \frac{c}{R^2} \cdot \sin(\theta)$$
 (D1)

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UNITED STATES PATENT AND TRADEMARK OFFICE  
CERTIFICATE OF CORRECTION

PATENT NO. : 4,993,416  
DATED : Feb. 19, 1991

INVENTOR(S) : Jonathan Ophir

It is certified that error appears in the above-identified patent and that said Letters Patent  
is hereby corrected as shown below:

Column 11, line 13, before "tissue" insert --ultrasonic--.

Signed and Sealed this

Fifteenth Day of December, 1992

Attest:

DOUGLAS B. CONER

Acting Commissioner of Patents and Trademarks

Attesting Officer

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